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Structural phase transitions in multipole traps

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(Dated: February 21, 2012)

A small number of laser-cooled ions trapped in a linear radiofrequency multipole trap forms a hollow tube structure. We have studied, by means of molecular dynamics simulations, the structural transition from a double ring to a single ring of ions. We show that the single-ring configuration has the advantage to inhibit the thermal transfer from the rf-excited radial components of the motion to the axial component, allowing to reach the Doppler limit temperature along the direction of the trap axis. Once cooled in this particular configuration, the ions experience an angular dependency of the confinement if the local adiabaticity parameter exceeds the empirical limit. Bunching of the ion structures can then be observed and an analytic expression is proposed to take into account for this behaviour.

I. INTRODUCTION

In the wide range of radiofrequency (rf) traps, quadrupole traps take a particular place due to the parabolic rf potential they create and to the harmonic pseudo-potential it generates. In contrast, higher order traps show a more "rectangle" shape of the rf electric field which induces a flat pseudo-potential well with a reduced rf-driven micromotion in the center of the trap [1]. This feature finds its application in the spectroscopy of cold molecular samples [2] or in the interrogation of the hyperfine transition of a large atomic sample for frequency metrology [3], the ions being cooled by collisions with a buffer gas.

Laser-cooling of ions trapped in multipole traps has only been performed very recently [4] and experimental observations have demonstrated a tube-like structure for the ion organization [5]. This self-organization in empty core structure results from the competition between Coulomb repulsion and a confining force very soft in the center of the trap and stiffer towards the walls. For large sample, this behavior is confirmed by the density profile deduced in the cold charged fluid limit [6] and for smaller ones, by simulations of the equilibrium structures [4, 7, 8]. For a large number of ions trapped in a linear multipole trap, the general structure of a crystallized sample is a set of concentric tubes for which the revolution axis is the longitudinal trap axis. The evolution of the structure with the number of ions and the trapping parameters is extensively studied in [8].

As an example, figure 1 shows the simulated structure of 1000 cold calcium ions inside the pseudo-potential of a linear octupole trap, laser-cooled to the Doppler limit temperature of 0.5 mK. For the same trapping parameters and a smaller number of ions, the structure reduces to a single tube, formed of parallel rings of ions aligned along the trap axis. For a decreasing number of ions, the tube reduces to two rings then to a single ring. Ring- or

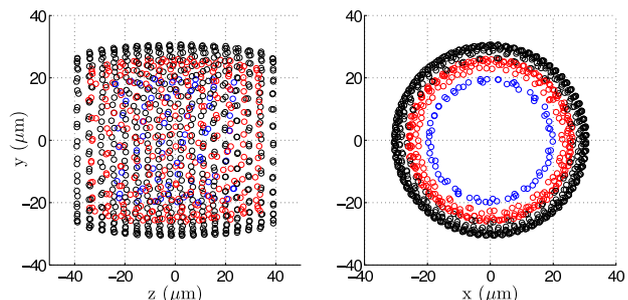


FIG. 1. (color online) Structure of a set of 1000 laser-cooled ions in the pseudo-potential of a linear octupole trap. The structure is composed of three concentric tubes extending along the trap axis, which is in the z direction. The temperature is 0.5 mK in the three directions of space.

tube-like structures offer the possibility to study dynamics in a two dimensional configuration. Furthermore, ion rings have symmetry properties that can be exploited in high-resolution spectroscopy. Among these highly symmetric structures, the single ring configuration has the outstanding property to be a true two dimensional system where the motional temperature along the symmetry axis can reach the Doppler limit when the ions are laser-cooled, despite the rf-driven motion in the transversal plane [9]. A previous numerical study [10] of the cooling properties of ion structures formed in quadrupole traps has shown that for structures of reduced spatial dimensions, Coulomb interaction does not always couple the different directions of motion of the ions. This decoupling inhibits the thermal transfer between the decoupled directions of the motion and allows to reach the Lamb-Dicke regime for the ion motion along the longitudinal trap axis [9].

In the present article we use molecular dynamics simulation of laser-cooled ions in a linear octupole trap to study their structural phase transition from a two-ring to a one-ring configuration. The model used to simulate the ion dynamics is described in section II together with some brief reminders on linear multipole traps. Section III presents an analysis of the structural transition

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itself and enlightens the role of the rf-driven motion on this transition. In section IV, we extend our study by increasing the adiabaticity parameter $\eta_{ad}(r)$ beyond the empirical limit.

II. FRAMEWORK OF THE MODEL

The numerical simulations use molecular dynamics, based on the symplectic velocity-Verlet algorithm to integrate Newton's equations. We have simulated the dynamics of a set of 100 ions in the potential of a linear octupole trap, cooled by Doppler laser-cooling. Ions are thus subject to conservative forces deriving from the trap electric potential and the Coulomb inter-ion potential, and dissipative forces due to Doppler cooling. The general form for the ideal rf potential of a linear $2k$ -pole trap can be written in polar coordinates like

$$\Phi_{2k}(r, \varphi, t) = q_e V_{2k} \left(\frac{r}{r_0} \right)^k \cos(k\varphi) \cos(\Omega t), \quad (1)$$

where V_{2k} is the amplitude of the rf potential applied to the $2k$ rods, oscillating at frequency $\Omega/2\pi$, r_0 is the inner radius of the trap, and q_e is the electric charge of the ions. Unless otherwise specified, we use the potential of an octupole trap ($2k = 8$) to run the simulations. This rf potential induces an rf-driven motion which produces a ponderomotive force, which leads to the radial confinement of the ions. It can be derived from the static pseudo-potential [1]:

$$\Psi_{2k}(r) = \frac{k^2}{4} \frac{q_e^2 V_{2k}^2}{m r_0^2 \Omega^2} \left(\frac{r}{r_0} \right)^{2k-2}. \quad (2)$$

The motion in this pseudo-potential has a time scale larger than an rf period and is called macro-motion.

The axial confinement is achieved by a harmonic static potential well characterized by the frequency $\omega_z/2\pi$, which scales like $\sqrt{q_e/m}$. Obeying Laplace's law, this static potential has a deconfining contribution in the radial plane. Hence, for each ion of mass m and position (r, φ, z) , the total trapping potential U_{trp} writes:

$$U_{\text{trp}}(r, \varphi, z, t) = \Phi_{2k}(r, \varphi, t) + \frac{1}{2} m \omega_z^2 \left(z^2 - \frac{r^2}{2} \right). \quad (3)$$

When the order of the multipole is higher than $2k = 4$, the deconfining contribution of the axial confinement leads to a shift of the minimum of the trapping potential in the radial plane, from the centre to a circle of radius R_{min} :

$$R_{\text{min}} = r_0 \left(\frac{1}{k^2 (k-1)} \frac{m r_0^2 \Omega^2}{q_e^2 V_{2k}^2} m r_0^2 \omega_z^2 \right)^{1/(2k-4)}. \quad (4)$$

Hence, contrary to a quadrupole trap configuration, the local minimum of the trapping potential is no more superposed to a node of the rf electric field [11]. Throughout

this article, the confinement parameters and number of ions are such that the ions sit very close to the potential minimum and they do undergo micro-motion.

The Coulomb interaction is taken into account by the total pair interaction potential, which writes:

$$U_c = \frac{q_e^2}{4\pi\epsilon_0} \sum_{i=1}^{N_p-1} \sum_{j=i+1}^{N_p} \frac{1}{|\delta_{ij}|}, \quad (5)$$

where N_p is the number of ions and δ_{ij} is the relative distance between ions i and j . The Doppler laser-cooling is modeled using velocity kicks simulating the ion's recoil, which happens depending on the probability laws of absorption (given by the stationary solution of the optical Bloch equations) and emission (given by the exponential decay law). For symmetry reasons, laser-cooling is applied in the three directions of the trap. Details about the model we used for the cooling are given in [10].

Initial conditions are prepared by giving to each ion an initial position randomly chosen in the inner 10% of the radius of the trap and a random velocity obeying a thermal distribution corresponding to 1 K. Ions are trapped by the rf electric field and are subject to Doppler cooling. The system is ready for further simulations when the temperatures are stationary. The temperature is one of the relevant parameters to get insight into ion dynamics. As the rf-driven motion is not a thermal motion, it has to be subtracted from the velocities before computing the temperature [12, 13]. In practice, this is done by averaging the velocity of the ions over an rf period. Each averaged component of the velocity is then taken into account to define the temperature associated to each direction of the motion. Explicit details about the computation of the temperatures can be found in [10].

All the forthcoming results assume perfect trap operation and neglect defects of the potential due to patch potential and/or misalignments of the electrodes as well as collisions with the residual gas. Patch potentials with a highly non-uniform spatial repartition could destroy the highly symmetric structure formed by few ions in a linear multipole trap. For high precision spectroscopy in multipole trap, more extra electrodes than for a line in quadrupole trap may be required to compensate for patch potentials and their gradient. Misalignment of the electrodes has been identified in a 22-pole trap [14] as responsible for 10 extra potential minima, which cause bunching of the ions in these local potential wells. As for collisions, they may perturb the formation process of the crystal but the trapping conditions chosen for these studies imply a strong confinement in the axial and in the radial direction, of the same order of magnitude as the one met in quadrupole trap. Therefore, the stationary structures described in this article should survive to collisions like ion chains do. This strong confinement may seem unrealistic regarding the smooth profile of the trapping field in the center of the trap. The scaling law for the rf electric field in an octupole is V_8/r_0^4 , and to reach a R_{min} value of $120\mu\text{m}$ with $\omega_z = 1$ MHz and $\Omega = 20$

MHz like chosen in the following, it takes an amplitude V_8 equal to 208 V in a 0.5 mm inner radius trap, which is very accessible. As for quadrupole trap dedicated to single ion experiment, a multipole trap able to keep a single ring stable has to be specifically designed to that purpose.

III. DOUBLE-RING TO SINGLE-RING STRUCTURAL TRANSITION

The trapping parameters are such that the Coulomb repulsion is negligible compared to the deconfining effect associated to the axial confinement and the radial position of the ions is controlled by the potential minimum R_{\min} . Depending on the number of ions N_p and the applied axial potential, the stationary structure is a single or a double ring. By stability consideration, one can show that there is a critical radius R_{crit} below which the equilibrium configuration is a double ring [6]. This critical radius only depends on N_p and ω_z through

$$R_{\text{crit}} \approx \left(\frac{q_e^2 / 4\pi\epsilon_0}{2 m \omega_z^2} \right)^{1/3} \frac{N_p}{\pi}, \quad (6)$$

For a set of 100 ions and $\omega_z/2\pi = 0.5$ MHz, the stability critical radius R_{crit} is 181 μm . The rf electric potential is chosen such that the radius R_{\min} is 120 μm and the ions organize themselves in two rings of 50 ions separated by an axial distance of about 9 μm . This double-ring structure can be seen on figure 2, for an rf phase corresponding to the maximal amplitude of the micro-motion. The

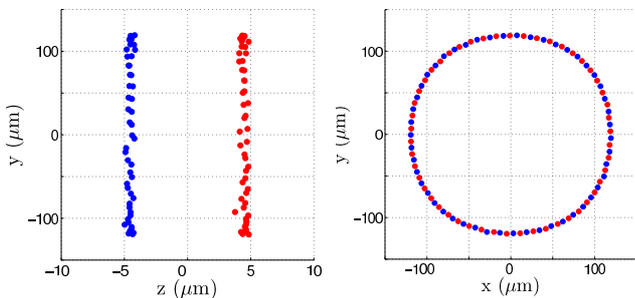


FIG. 2. (color online) Double-ring structure of 100 laser cooled ions, in the rf potential of an octupole trap. The axial confinement frequency is $\omega_z/2\pi = 0.5$ MHz, leading to an axial separation between the two rings of about 9 μm . The picture corresponds to the instant of the maximal amplitude of the micro-motion.

pattern formed by the two ion rings is equivalent to an annular zig-zag structure [15], the positions of two consecutive ions being on different sides of the $z = 0$ plan. This perfect annular zig-zag is obtained by choosing an even number of ions. In this configuration, the temperature associated with the axial confinement of the motion is $T_z \approx 1$ mK (see figure 3). If the velocity of the ions is expressed in cartesian coordinates, the T_x and T_y temperatures are identical and are approximately equal to 12 mK.

The difference between these values is attributed to a partial thermalization of the degrees of motion, the higher temperature being induced by rf-driven motion in the radial plane. This motion induces rf-heating by which the kinetic energy of the rf-driven motion is transferred to the macro-motion [16]. It also reduces the Doppler cooling efficiency by the Doppler shift on the cooling transition. The temperature discrepancy shows that the Coulomb coupling in the two-ring structure is not strong enough to thermalize the motion in the radial plane with the motion along the axis [10]. Inside the radial plane itself, the use of cylindrical coordinates reveals an important discrepancy in the temperature associated to the radial motions ($T_r \simeq 10$ mK) and the one associated to the tangential motion ($T_\varphi \simeq 2$ mK, see figure 3).

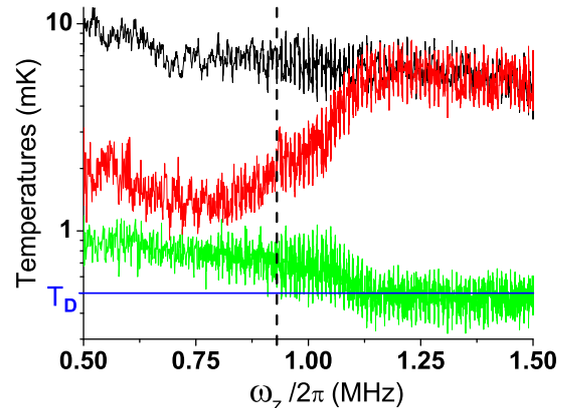


FIG. 3. (color online) Temperatures associated to the motion in a polar coordinate system T_r (black upper curve), T_φ (red medium curve) and T_z (green bottom curve), when the axial confinement frequency ω_z is increased. The vertical dashed line marks the theoretical transition frequency from the double-ring to the single-ring structure. The horizontal blue line materializes the Doppler limit temperature.

Starting from this configuration, we study the double to single ring transition. To that purpose, the characteristic frequency of the axial confinement $\omega_z/2\pi$, is slowly increased from 0.5 MHz to 1.5 MHz. At the same time, the amplitude of the rf electric field is raised to balance the enhancement of the static deconfining effect in the radial plane, keeping the minimum of the trapping potential, R_{\min} , and thus the radius of the ring, constant (see equation 4). For a constant number of ions, the structural transition is expected from equation 6 to take place for $\omega_z/2\pi = 0.93$ MHz. It is verified that the variation of the trapping parameters is sufficiently slow to consider the system to be in a stationary state during the simulation. Particular cares are taken for values above 0.92 MHz for which the increase of the confinement parameters is halted every 0.1 MHz, by time periods of 10 ms.

The structural transition is first monitored by the distance between the center of mass (c.m.) of the two rings, Δz_{cm} , given on the left axis of figure 4, as a function of

the characteristic frequency $\omega_z/2\pi$. The value for which the two rings merge is in good agreement with the theoretical frequency of 0.93 MHz, for which the structural transition is predicted.

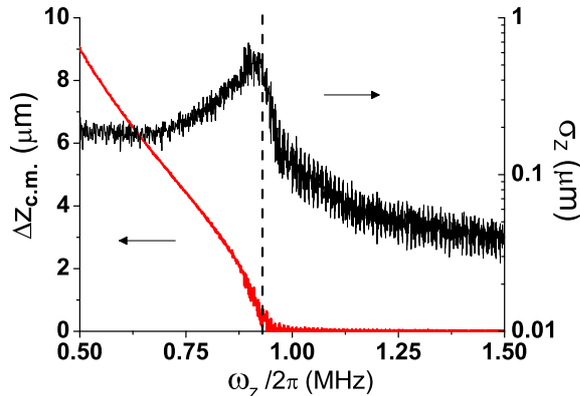


FIG. 4. (color online) Relative distance between the c.m. of the two rings, Δz_{cm} (red curve, left axis), and standard deviation of the axial component of the ion's positions from the c.m. position of the ring of which they belongs, σ_z (black curve, right axis), function of the increasing axial confinement frequency ω_z . The vertical dashed line marks the theoretical transition from the double-ring to single-ring configuration.

The standard deviation of the axial component of the ions' positions from the c.m. position is also plotted on figure 4. For $\omega_z/2\pi = 0.5$ MHz, the standard deviation σ_z is due to the axial thermal motion around the equilibrium position. Increasing ω_z , the axial temperature (shown on figure 3) drops while the local steepness of the potential in the axial direction is increased. Hence, the axial dispersion on ions' positions which is due to thermal motion is expected to decrease when ω_z grows. Figure 4 shows that it becomes maximum around the structural transition before decreasing to a lower value than the initial one. To understand the transition process, we focus on the ion organization just before the structural transition. While the two rings are getting closer, a distortion of the ring structures progressively appears, visible on figure 5 and stationary over the rf period. The distortion does not affect the equilibrium radius of the structures but the axial inter-ion distance is smaller for ions localized close to the rf electrodes than for those which are trapped far from the electrodes (with the convention defined by equation 1, electrodes are at angular positions $\varphi_n = n\pi/4$). This behaviour is attributed to micro-motion whose amplitude $\vec{\varepsilon}_\mu(r, \varphi, t)$ has the analytic form:

$$\vec{\varepsilon}_\mu(r, \varphi, t) = \frac{q_e V_8 r^3}{m r_0^4 \Omega^2} (\cos(4\varphi) \hat{r} - \sin(4\varphi) \hat{\varphi}) \cos(\Omega t), \quad (7)$$

When ions are localized in front of an rf-electrode, the micro-motion amplitude is almost exclusively radial, and has no significant consequence on the relative distance of two neighbouring ions belonging to different rings. On

the contrary, when ions are localized at an inter-electrode angular position, the micro-motion is mostly tangential to the ring structure and depends like $\sin(4\varphi)$ on the ion angular position. Two neighbouring ions, separated by an angle $2\pi R/N_p$, do not have the same tangential amplitude of micro-motion, producing an oscillation of their angular separation. Since the neighbouring ions do not belong to the same ring, this variation of angular positions brings them face to face, increasing their Coulomb repulsion in the axial direction. The comparison with simulations run in the pseudo-potential (equation 2) confirms that this behaviour is induced by the rf motion.

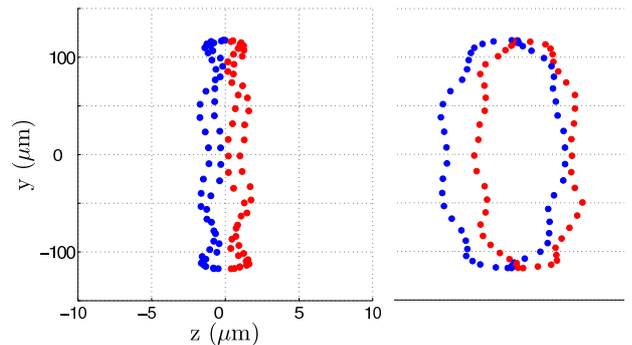


FIG. 5. (color online) Distorsion of the double-ring structure viewed in the (z, y) -plane (left picture), and from a point of view rotated of $-\pi/4$ around the y -axis. The structure is stationary over the rf-period time scale. This picture is taken when the distance between the two centers of mass is about $2 \mu\text{m}$.

The evolution of the distortion during the increase of ω_z explains the increase of standard deviation σ_z roughly situated between 0.75 MHz and 0.93 MHz (Figure 4). When the transition frequency is reached, the c.m. of the two structures match roughly, but the standard deviation takes its maximal value, showing that the structure is not yet a planar ring. For higher values of ω_z , the standard deviation shows a steep downward slope, until a frequency value slightly lower than 1 MHz. The structure can then be considered as a planar single-ring structure.

The distortion of the structure described above and the coupling between the tangential and axial component of motion it induces, explain the initial discrepancy between the radial and tangential temperatures when the structure is a double-ring (see figure 3). The axial motion being not subjected to micro-motion, it can reach lower temperature than the motions in the radial plane. The coupling between axial and tangential components of the motion allows to dissipate a part of the tangential thermal kinetic energy through the axial cooling. As the frequency ω_z is increased, the temperature T_φ increases progressively and meets the temperature associated to the radial directions, T_r . At the same time, the axial temperature T_z shows a steeper drop between 1 MHz

and 1.25 MHz, reaching the Doppler limit temperature at the frequency where the two other temperatures join. This behaviour suggests that the axial component of the motion of the ions is then decoupled from the two transverse directions, like expected for a full two dimensional system [10].

The comparison between Figures 4 and 3 shows that for this small 100-ion system, informations on the dimensionality of the system given by the variation of the ions' positions, σ_z , and the temperatures of the system do not coincide. The decoupling observed on the temperature arises when the σ_z parameter already shows a planar structure. As a consequence, each of these parameters may be a relevant criteria to determine the structural phase transition.

IV. ABOVE THE ADIABATIC LIMIT

The single ring structure presented in the preceding section seems very promising for applications such high precision spectroscopy, quantum information or for metrological applications [9, 17]. For all of these, it is mandatory to reduce the amplitude of motion of the ions to below the chosen transition wavelength (the Lamb-Dicke regime [18]) to cancel the first order Doppler effect. For trapped ions, Doppler laser-cooling is sufficient if the trapping frequency along the laser propagation direction is large enough. In practice, for calcium ions cooled to the Doppler limit, a motional frequency of 1 MHz allows the ions to reach the Lamb-Dicke regime for an optical transition. One of the main drawbacks of the ring structure compared to its linear counterpart, the ion chain, found in quadrupole traps is the non null rf-driven motion (micro-motion) at their equilibrium positions. Figure 3 shows that in the single ring configuration, the motion along the axis is decoupled from the motion in the radial plane where micro-motion, and thus rf-heating, take place. The Lamb-Dicke regime can thus be reached for laser propagating along the trap axis.

To quantify the impact of the micro-motion on the ion dynamics and laser cooling, the Mathieu parameter q_x , relevant for this issue in quadrupole traps, can be extrapolated to multipole trap and is then related to the more general concept of adiabaticity of the trajectories [1]. The adiabaticity parameter η_{ad} depends on the radial position of the ions inside the trap and at the potential minimum R_{\min} , it takes the simple expression [6]:

$$\eta_{ad}(R_{\min}) = 2 \sqrt{k-1} \omega_z / \Omega . \quad (8)$$

It has been demonstrated, numerically and experimentally in a 22-pole [2, 19, 20], that for $\eta_{ad}(r) < 0.36$ the trajectories are stable but few studies concern this adiabaticity parameter in the context of cold samples. In this section, we focus on the role of the micro-motion on the self-organization of the ions and relate it to the adiabaticity parameter. Figure 6 shows the single ring structure formed by 100 ions located at the same radial

potential minimum and trapped by an axial potential characterized by $\omega_z/2\pi$ equal to 1 MHz ($\eta_{ad} = 0.173$), 2 MHz ($\eta_{ad} = 0.346$) and 3 MHz ($\eta_{ad} = 0.520$). For increasing ω_z , we observe a bunching of the ions at the inter-electrode angular positions.

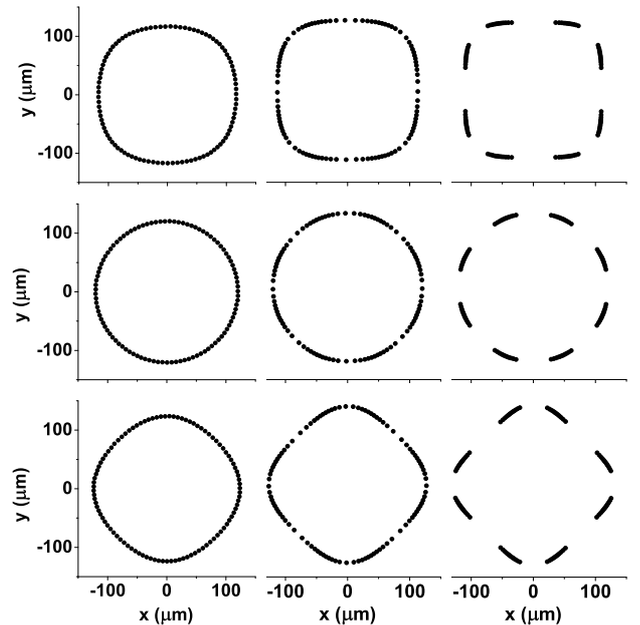


FIG. 6. View in the radial plane of structures obtained for an equilibrium radius $R_{\min} = 120 \mu\text{m}$ and, from left to right, $\omega_z = 1$ MHz, 2 MHz and 3 MHz. From top to bottom, the pictures correspond to three phase of the rf: $\cos(\Omega t) = -1; 0; 1$.

This bunching is not observed in the pseudo-potential approach of the trapping electric field and is attributed to the micro-motion of the ions. Some insight in the role of micro-motion can be gained by considering a cold sample sitting close to R_{\min} , where the radial trapping potential can be approximated by a local harmonic potential. Assuming a null potential for R_{\min} , it can be written like

$$U_{1oc}(R_{\min} + \delta r) = \frac{1}{2} m \omega_{1oc}^2 (\delta r)^2 , \quad (9)$$

with the characteristic frequency $\omega_{1oc} = \sqrt{k-2} \omega_z$ [6]. The ion motion is assumed to be due only to the rf-driven micro-motion whose amplitude can be written like:

$$\vec{\varepsilon}_{\mu}(R_{\min}, \varphi, t) = \|\vec{\varepsilon}_{\mu}(R_{\min})\| (\cos(4\varphi) \hat{r} - \sin(4\varphi) \hat{\varphi}) \cos(\Omega t) , \quad (10)$$

where the amplitude of the micro-motion $\|\vec{\varepsilon}_{\mu}(R_{\min})\|$ is easily deduced from equation 7. Only the radial contribution of this motion engenders a significant potential energy variation during the rf-period. Its time average depends on the angular position φ like

$$\overline{U_{1oc}}(R_{\min}, \varphi) = \frac{1}{4} m \omega_{1oc}^2 \|\vec{\varepsilon}_{\mu}(R_{\min})\|^2 \cos^2(4\varphi) \quad (11)$$

and could explain the bunching observed on figure 6. To test this assumption, we have created a configuration

where only two ions bunch in one of the local angular minimum potential localized around $\varphi_0 = (2n + 1)\pi/8$. To that purpose, we have simulated a set of 9 ions in the rf-potential of the same octupole trap, with $\omega_z/2\pi$ initially set at 0.5 MHz, and slowly increased to 4 MHz, keeping $R_{\min} = 120 \mu\text{m}$ constant by increasing the rf potential. The final stable configuration is then made of seven ions and a unique pair of ions, localized at each inter-electrode angular positions.

Assuming a local description $-m\omega_{\text{mes}}^2\delta/2$ for the force responsible for the bunching, the measure of the relative distance δ of the ions forming the pair allows one to deduce the characteristic frequency ω_{mes} from the equilibrium condition with the Coulomb repulsion:

$$\omega_{\text{mes}} = \left(\frac{2 q_e^2}{4\pi\epsilon_0 m \delta^3} \right)^{1/2}. \quad (12)$$

One can also evaluate the force \vec{F}_φ by deriving the time-averaged micro-motion induced potential $\overline{U}_{\text{loc}}(R_{\min}, \varphi)$, for an angle $\varphi = \varphi_0 + \delta\varphi$:

$$\vec{F}_\varphi(\varphi_0 + \delta\varphi) = -8m\omega_{\text{loc}}^2 \frac{\|\vec{\varepsilon}_\mu(R_{\min})\|^2}{R_{\min}} \delta\varphi \hat{\varphi} \quad (13)$$

Writing \vec{F}_φ like $-m\omega_\varphi^2 R_{\min} \delta\varphi$, one can connect the characteristic frequency ω_φ to the local radial potential. The results of numerical simulations show that the predicted bunching frequency ω_φ is larger than the measured frequency ω_{mes} by a factor $\sqrt{2}$, all along the simulation increasing ω_z at constant R_{\min} . The disagreement by a factor $\sqrt{2}$ is also observed when the same simulations are run with 13 ions in a dodecapole trap. According to the results obtained from the simulations, the local bunching characteristic frequency is given by:

$$\omega_\varphi = 2 \omega_{\text{loc}} \frac{\|\vec{\varepsilon}_\mu(R_{\min})\|}{R_{\min}}. \quad (14)$$

Eq. 14 shows how the bunching frequency depends on the characteristic trapping parameter and ring configuration. Indeed, using the dependence of the micro-motion with the position in the trap, one can show that ω_φ scales like ω_z^2/Ω . Hence, the bunching force scales like $R_{\min}\omega_z^4/\Omega^2$. The dependence of this bunching force with ω_z was illustrated on figure 6 for fixed R_{\min} . Its dependence with R_{\min} for fixed ω_z can be seen on figure 7 and confirms an increase of the bunching with R_{\min} . These different studies show that the so-called adiabaticity parameter η_{ad} is not the relevant criteria to estimate the role of micro-motion on the ion organization. Indeed, the four structures plotted on figure 7 share the same η_{ad} . This behaviour shows that in the general description of the ion dynamics in a multipole trap, micro-motion has a non-negligible influence not only on the thermal properties of the sample but also on its structural properties, and has thus to be taken into account.

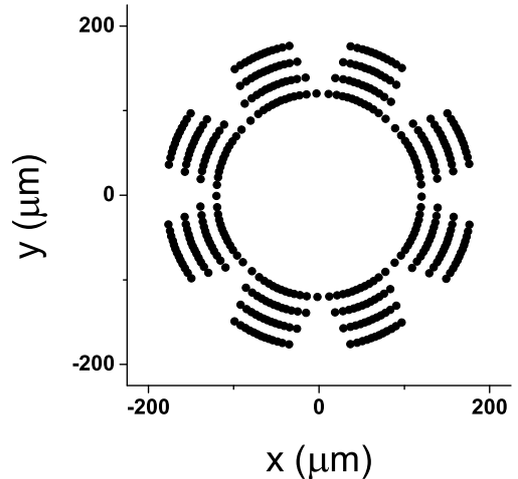


FIG. 7. View in the radial plane of a 100-ion structure, for $\omega_z/2\pi = 2$ MHz, and $R_{\min} = 120 \mu\text{m}$, $140 \mu\text{m}$, $160 \mu\text{m}$ and $180 \mu\text{m}$, varied by modifying the rf-potential amplitude. The picture corresponds to a null rf-phase.

V. CONCLUSION

In this paper, we have numerically studied the doubling to single-ring structural transition of a Doppler-cooled 100-ion system in the rf potential of an octupole trap. The simulations showed that, in the double-ring configuration, the temperatures associated to each component of the motion expressed in cylindrical coordinates are different. In our initial configuration, the tangential temperature is lower than the radial one by almost one order of magnitude. This is attributed to a coupling of the tangential motion to the axial one, which is efficiently Doppler-cooled due to the absence of rf-driven motion along the axial direction. When the double-ring structure is compressed, the c.m. of each ring merge at the predicted transition frequency but the structure shows a distortion which is maximal at this frequency. This behaviour is attributed to the micro-motion of the ions, inducing an axial Coulomb repulsion which can be considered as a pseudo-potential between the ions of the different rings. A planar single-ring structure is thus obtained for a higher value of the predicted transition frequency. When ω_z is further increased, the axial temperature reaches the Doppler limit value showing a decoupling of the axial and transverse components of the motion.

For a cold ion system, we have also shown that it is possible to increase the adiabaticity parameter beyond the empirical limit, keeping stable ion confinement. At a fixed ring radius, increasing the adiabaticity parameter results in a bunching of the structure, for which an expression of the local bunching frequency is proposed. We have also shown that changing the ring radius at constant value of the adiabaticity parameter leads to the

emergence of the bunching phenomenon.

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